EXP GANGLIOSIDE GD3 EXP GANGLIOSIDE GD3/CN 2 S E3 FILE 'HCAPLUS' ENTERED AT 13:08:49 ON 30 APR 2010 121 S L1/THU

FILE 'REGISTRY' ENTERED AT 13:08:16 ON 30 APR 2010

L2

L3 390905 S INFLAMM? OR ANTIINFLAMM?

L1

L4 20 S L2 AND L3 L5 265180 S CHOLESTEROL OR HYPERCHOLESTEROLEM? OR ATHEROSCLEROSIS

L6 8 S L2 AND L5 => file registry COST IN U.S. DOLLARS

SINCE FILE

TOTAL. ENTRY SESSION 0.22 0.22

FILL ESTIMATED COST

FILE 'REGISTRY' ENTERED AT 13:08:16 ON 30 APR 2010 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2010 American Chemical Society (ACS)

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 29 APR 2010 HIGHEST RN 1220951-91-6 DICTIONARY FILE UPDATES: 29 APR 2010 HIGHEST RN 1220951-91-6

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TSCA INFORMATION NOW CURRENT THROUGH January 8, 2010.

Please note that search-term pricing does apply when conducting SmartSELECT searches.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/support/stngen/stndoc/properties.html

```
=> exp ganglioside GD3
                3
                       GANGLIOSID/BI
E1
E2
             1271
                        GANGLIOSIDE/BI
E3
                0 --> GANGLIOSIDE GD3/BI
E4
                       GANGLIOSIDES/BI
               GANGLIOSIDES/BH
GANGLIOSIDOL/BI
GANGLIOSIDOSIS/BI
GANGLIOSPHINGO/BI
GANGLIOSPHINGOSINE/BI
GANGLIOSTAT/BI
GANGLIOTETRA/BI
GANGLIOTETRA/BI
GANGLIOTETRAOSE/BI
E5
E6
E7
E8
E9
E10
E11
E12
                3
                        GANGLIOTETRAOSYL/BI
=> exp ganglioside GD3/cn
E1
                        GANGLIOSIDE GD2, N'-ACETYL-N-GLYCOLOYL-/CN
                 1
E2
                         GANGLIOSIDE GD2, N.N'-DIACETYL-/CN
E3
                 2 --> GANGLIOSIDE GD3/CN
E4
                        GANGLIOSIDE GD3 (SYNTHETIC)/CN
                 1
E5
                         GANGLIOSIDE GD3 B1,4-N-ACETYLGALACTOSAMINYLTRANSFERASE/
                 1
                         CN
                      GANGLIOSIDE GD3 ACETYLGALACTOSAMINYLTRANSFERASE/CN
GANGLIOSIDE GD3 AMIDE/CN
GANGLIOSIDE GD3 LACTONE I/CN
GANGLIOSIDE GD3 LACTONE II/CN
GANGLIOSIDE GD3 SYNTHASE/CN
E6
                1
E7
                 1
E8
                1
E9
                 1
                1
E10
E11
                1
                       GANGLIOSIDE GD3 SYNTHASE (HUMAN CLONE PAMO-GD3)/CN
E12
                1
                        GANGLIOSIDE GD3 SYNTHETASE/CN
```

=> s e3

2 "GANGLIOSIDE GD3"/CN

=> d 11 scan

- L1 2 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN Ceramide,  $1-0-[0-(N-acetyl-\alpha-neuraminosyl)-(2+8)-0-(N-acetyl-\alpha-neuraminosyl)-(2+3)-0-<math>\beta$ -D-galactopyranosyl-(1+4)- $\beta$ -D-galucopyranosyl]-
- MF Unspecified
- CI COM, MAN
- \*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*
- \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

HOW MANY MORE ANSWERS DO YOU WISH TO SCAN? (1):1

- L1 2 ANSWERS REGISTRY COPYRIGHT 2010 ACS on STN
- IN Ganglioside GD3
- MF Unspecified
- CI COM, MAN
- \*\*\* STRUCTURE DIAGRAM IS NOT AVAILABLE \*\*\*
- \*\*PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT\*\*

ALL ANSWERS HAVE BEEN SCANNED

=> file hcaplus COST IN U.S. DOLLARS FULL ESTIMATED COST

SINCE FILE TOTAL ENTRY SESSION 5.99 6.21

FILE 'HCAPLUS' ENTERED AT 13:08:49 ON 30 APR 2010 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2010 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 30 Apr 2010 VOL 152 ISS 19
FILE LAST UPDATED: 29 Apr 2010 (20100429/ED)
REVISED CLASS FIELDS (/NCL) LAST RELOADED: Feb 2010
USPTO MANDAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Feb 2010

HCAplus now includes complete International Patent Classification (IPC) reclassification data for the first quarter of 2010.

CAS Information Use Policies apply and are available at:

http://www.cas.org/legal/infopolicy.html

This file contains CAS Registry Numbers for easy and accurate substance identification.

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=> s 11/t.hu
          1576 L1
       1237249 THU/RL
L2
           121 L1/THU
                 (L1 (L) THU/RL)
=> s inflamm? or antiinflamm?
        381286 INFLAMM?
         66629 ANTIINFLAMM?
1.3
        390905 INFLAMM? OR ANTIINFLAMM?
=> s 12 and 13
L4
           20 L2 AND L3
=> d 14 1-20 ti abs bib
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- ANSWER 1 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN
- TI Anti-clucan antibody recruitment of complement-mediated immune responses
- AB The authors disclose the presence of human natural antibodies to β1-6-glucans. In one example, anti-β1-6 glucan IgG antibodies are shown to mediate complement activation and neutrophil phagocytosis. In a second example, a conjugate of Herceptin with β1-6 glucan was shown to target breast cancer cells for complement-mediated lysis and recruitment of neutrophils.
- AN 2009:1366143 HCAPLUS <<LOGINID::20100430>>
- DN 151:526825
- ΤI Anti-glucan antibody recruitment of complement-mediated immune responses
- Rubin-Bejerano, Ifat; Fink, Gerald R.; Kohane, Daniel S. IN
- PA Immunexcite, Inc., USA
- PCT Int. Appl., 74 pp. SO CODEN: PIXXD2
- DT Patent
- LA English

FAN.	CNT	1																	
	PAT	TENT :	NO.			KIN	D	DATE			APPL	ICAT	I NOI	NO.		D.	ATE		
							_									-			
PI	WO	2009	1348	91		A2		2009	1105		WO 2	009-	JS42	117		2	0090	429	
	WO	2009	1348	91		A3		2010	0218										
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			FI,	GB,	GD,	GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	
			KG,	KM,	KN,	KP,	KR,	KZ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LY,	MA,	MD,	
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			TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	zw			
		RW:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HR,	HU,	
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	ZW, AM, AZ					BY,	KG,	KZ,	MD,	RU,	ΤJ,	TM,	ΑP,	EA,	EP,	OA			
PRAT	IIS	2008	-714	37P		P		2008	0429										

PRAI US 2008-71437P

- T. 4 ANSWER 2 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN
- ΤТ Compositions comprising phospholipids
- The present invention provides compns. comprising phospholipids and particularly those comprising at least 40% phospholipid and at least 80%

phospholipid as a percentage of total fat in the extract, comprising polyunsatd. and saturated phospholipids, in a ratio of saturated phospholipid

monounsatd. to polyunsatd. phospholipid of about 6:3:1 resp., or comprising at least 40% phospholipid and less than 40% protein and methods for their production from dairy products.

AN 2009:239236 HCAPLUS <<LOGINID::20100430>>

DN 150:258888

tο

I Compositions comprising phospholipids

IN Brown, Andrew; Rowney, Michelle

PA Murray Goulburn Co-Operative Co. Limited, Australia

SO PCT Int. Appl., 80pp.

CODEN: PIXXD2

DT Patent

LA English FAN.CNT 1

	PAT	ENT :				KIN		DATE			APPL						ATE	
PI		2009															0080	
		W:	ΑE,	AG,	AL,	AM,	AO,	AT,	AU,	AZ,	BA,	BB,	BG,	BH,	BR,	BW,	BY,	BZ,
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			FI,	GB,	GD,	GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,
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			ME,	MG,	MK,	MN,	MW,	MX,	MY,	ΜZ,	NA,	NG,	ΝI,	NO,	ΝZ,	OM,	PG,	PH,
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		RW:						CZ,										
								LV,										
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								MD,										
		2008																
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		R:						CZ,										
								LU,	LV,	MC,	MT,	NL,	NO,	PL,	PT,	RO,	SE,	SI,
						BA,												
PRAI		2007																
		2008																
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			AL:	r CI	TATI	ONS .	AVAI	LABL	E IN	THE	RE I	FORM	AT					

- L4 ANSWER 3 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN
- TI Method and apparatus of low strength electric field network-mediated delivery of drug, gene, siRNA, shRNA, protein, peptide, antibody or other biomedical and therapeutic molecules and reagents in skin, soft tissue, joints and bone
- AB The invention includes four preferred embodiments: (i) a method and apparatus for the joint and its related soft tissue for bone gene, protein and drug delivery; (ii) a method and apparatus for gene, protein and drug delivery to its method and apparatus for delivery of gene, protein and drug delivery to skin and soft tissue; and/or (iv) a method and apparatus for delivery of a gene, protein and drug to soft tissue tumor. The apparatus for transfecting drug, gene, siRNA, shRNA, protein, peptide, antibody or other biomedical and therapeutic mols. and reagents comprises a plurality of neg. electrodes disposed into low resistance elec. contact with skin overlaving the tissue.
- AN 2007:1207931 HCAPLUS <<LOGINID::20100430>>
- DN 147:474740
  - Method and apparatus of low strength electric field network-mediated delivery of drug, gene, siRNA, shRNA, protein, peptide, antibody or other biomedical and therapeutic molecules and reagents in skin, soft tissue,

```
joints and bone
TN
    Sen, Luyi
    The University of California, USA
PA
SO
   PCT Int. Appl., 51 pp.
    CODEN: PIXXD2
    Pat.ent.
LA.
   English
FAN.CNT 1
                    KIND DATE APPLICATION NO. DATE
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    WO 2007120557 A2 20071025 WO 2007-US8445
WO 2007120557 A3 20081113
                                                                    20070402
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             CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB,
             GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM,
             KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME, MG,
             MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT,
             RO, RS, RU, SC, SD, SE, SG, SK, SL, SM, SV, SY, TJ, TM, TN, TR,
             TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW
         RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
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         2647520 Al 20071025 CA 2007-2647520 20070402

8001519 A2 20081217 EP 2007-774731 20070402

R: AT, BE, BG, CH, CT, CE, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE,
     CA 2647520
     EP 2001519
             IS, IT, LI, LT, LU, LV, MC, MT, NL, PL, PT, RO, SE, SI, SK, TR,
             AL, BA, HR, MK, RS
                                                                    20070830
     CN 101506370 A 20090812 CN 2007-80000069
                             20090812
20090320
20060410
20060706
                         A
     IN 2008CN05422
                                            IN 2008-CN5422
                                                                     20081010
PRAI US 2006-744528P P WO 2007-US8445 W
                                20070402
OSC.G
              THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)
L4
     ANSWER 4 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN
ΤI
     Formulations for mediating inflammatory bowel disorders
AB
    The invention provides formulations and methods for mediating
     inflammation, in particular an inflammatory bowel
     disorder such as necrotizing enterocolitis. Further, the formulations are
     effective in lowering blood cholesterol and decreasing blood cholesterol
     absorption. The formulations comprise at least one ganglioside, which may
     be selected from the group consisting of: GD3, GM1, GM2, GM3, and GD1b.
     The invention provides a method of treating or preventing
     inflammatory diseases, such as necrotizing enterocolitis by
     delivery of at least one ganglioside to a subject in need thereof.
     Supplementation of foods or liqs. with gangliosides, for example infant
     formula or infant foods, can be employed according to the invention.
    2007:815148 HCAPLUS <<LOGINID::20100430>>
AN
    147:197354
DN
ΤI
    Formulations for mediating inflammatory bowel disorders
IN
    Clandinin, Michael Thomas; Park, Eek J.
    Mti Meta Tech Inc., Can.
    U.S. Pat. Appl. Publ., 39pp., Cont.-in-part of U.S. Ser. No. 551,789
     CODEN: USXXCO
    Patent
LA English
FAN.CNT 2
     PATENT NO. KIND DATE APPLICATION NO. DATE
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PΤ
    US 20070173480
                        A1
                               20070726 US 2007-622858
                                                                  20070112
    WO 2004087173
                         A2
                               20041014
                                          WO 2004-CA375
                                                                  20040312
    WO 2004087173
                         A3
                               20041125
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
            CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD,
            GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC,
            LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI,
            NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY,
            TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW
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            SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN,
            TD, TG
    US 20060276430
                         A1
                               20061207
                                          US 2004-551789
                                                                  20040312
PRAI US 2004-551789
                         A2
                               20040312
    WO 2004-CA375
                               20040312
                         Ta7
    US 2003-404095
                         Α
                               20030402
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

- ANSWER 5 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN
- Gangliosides as binding agents for vacuolating toxin VacA of Helicobacter pylori, their use for drugs and for foods, and screening method for therapeutic or prophylactic drugs for H. pylori-related diseases
- The binding agents for VacA, vacuolating toxin of H. pylori, contain AB gangliosides, lysogangliosides, and/or their chemical modification products. VacA activity is neutralized by the VacA-binding agents for treatment or prevention of H. pylori-related diseases, e.g., gastritis, gastric ulcer, and gastric cancer. The VacA-binding agents or VacA are used for screening of therapeutic or prophylactic drugs for H. pylori-related diseases. Gangliosides, lysogangliosides, and/or their chemical modification products are used for manufacture of therapeutic or prophylactic drugs for H. pylori-related diseases and for foods for suppression of the actions of H. pylori. Ganglioside GM1 (at 50 µg/mL) significantly inhibited the vacuolating activity of VacA in cultured human gastric epithelial cancer cell line AZ-521.
- AN 2007:251881 HCAPLUS <<LOGINID::20100430>>
- DN 146:266771
- TΙ Gangliosides as binding agents for vacuolating toxin VacA of Helicobacter pylori, their use for drugs and for foods, and screening method for therapeutic or prophylactic drugs for H. pylori-related diseases
- IN Wada, Akihiro; Hirayama, Toshiya; Yamazaki, Shigeki; Maeda, Kayo; Hasegawa, Makoto
- PA Nagasaki University, Japan; Kansai Bunri Sougougakuen
- SO Jpn. Kokai Tokkyo Koho, 14pp.
- CODEN: JKXXAF Patent DT
- LA Japanese
- FAN.CNT 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI JP 2007055921 PRAI JP 2005-241839	A	20070308	JP 2005-241839	20050823
FRAI OF 2000-241009		20030023		

- ANSWER 6 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN T. 4
- The immune response to disialoganglioside GD3 vaccination in normal dogs: a melanoma surface antigen vaccine
- As a result of its metastatic potential, canine malignant melanoma like its human counterpart like its human counter part, has a poor response to conventional treatment protocols. This prompted us to investigate the possibility of enhancing the immune response against the melanoma cell

surface antigen, disialoganglioside GD3. Initially a flow cytometric study was designed in which the incidence of GD3 on the cell surface, recognized by the monoclonal antibody Mel-1 (R24), was established in canine melanoma cell lines. Results from the flow cytometry found GD3 to be highly expressed (94.2%) in six out of seven canine melanoma cell lines. Since it was thus potentially a good target, a study in which normal dogs were vaccinated intradermally with a vaccine containing GD3 plus adjuvants was designed. The adjuvant included CpG oligodeoxynucleotide (CpG-ODN) sequences and RIBI-adjuvant, which are known to target toll-like receptors (TLR) of the innate immune system. From a cohort of 10 dogs, 4 were vaccinated 3 times, at 4 weekly intervals with GD3 plus adjuvant, and 4 received only RIBI-adjuvant, and 2 phosphate buffered saline. Caliper measurements were collected to assess skin reaction at the vaccination site and sera assayed for IqM and IqG antibodies against GD3 and cell-mediated cytotoxicity against a melanoma cell line. Results from the study found significant differences (P < 0.05) in the vaccine site reactions, IgM/IgG levels and cell-mediated cytotoxicity in the vaccinated vs. unvaccinated dogs. The addition of CpG-ODN sequences and increasing GD3 concentration in the vaccine increased the inflammation response at the injection site. GD3 IqG and IqM antibodies in vaccinated dogs showed increasing titers over time and achieved significance at weeks 9 and 12, resp. Cell-mediated cytotoxicity was only detected in peripheral blood mononuclear cells from vaccinated dogs. In conclusion, by combining the tumor antigen GD3 (a known weak self-antigen) and an adjuvant, tolerance was overcome by an innate and adaptive immune response in this population of normal dogs.

AN 2006:1139866 HCAPLUS <<LOGINID::20100430>>

DN 146:226964

TI The immune response to disialoganglioside GD3 vaccination in normal dogs: a melanoma surface antigen vaccine

AU Milner, R. J.; Salute, M.; Crawford, C.; Abbot, J. R.; Farese, J.

CS Department of Small Animal Clinical Sciences, College of Veterinary Medicine, University of Florida, Gainesville, FL, USA

SO Veterinary Immunology and Immunopathology (2006), 114(3-4), 273-284 CODEN: VIIMDS; ISSN: 0165-2427

PB Elsevier B.V.

DT Journal

LA English

RE.CNT 62 THERE ARE 62 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 7 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN

TI Bispecific antibodies, chimeric antibodies and fragments specific to human CD3 complex and antigen or tumor antigen for diagnosis and treatment of cancer, inflammation, allergy, infection and autoimmune disease

The present invention provides a bispecific binding mol., wherein said AB mol. comprises or consists of at least two domains whereby one of said at least two domains specifically binds to/interacts with the human CD3 complex and said domain comprises an amino acid sequence of an antibody derived light chain, wherein said amino acid sequence is a particularly identified amino acid sequence comprising specific amino acid substitutions, and a second domain is or contains at least one further antigen-interaction-site and/or at least one further effector domain. The invention further provides nucleic acid mols. encoding the bispecific binding mols. of the invention, vectors comprising said nucleic acid mols. and host cells transformed or transfected with said vectors. Moreover, the invention concerns a method for the production of bispecific binding mols. of the invention and compns. comprising the bispecific binding mols. of the invention, the nucleic acid mols. of the invention or the host cells of the invention.

AN 2005:902921 HCAPLUS <<LOGINID::20100430>>

- DN 143:246762
- II Bispecific antibodies, chimeric antibodies and fragments specific to human CD3 complex and antigen or tumor antigen for diagnosis and treatment of cancer, inflammation, allergy, infection and autoimmune disease
- IN Kufer, Peter; Lenkkeri-Schuetz, Ulla; Lutterbuese, Ralf; Kohleisen, Birgit PA Micromet A.-G., Germany
- SO PCT Int. Appl., 94 pp.
- CODEN: PIXXD2
- DT Patent
- LA English FAN.CNT 1

PATENT NO.	KI	ND	DATE			APPL	ICAT	ION I	NO.		D	ATE	
PI WO 2005077982	A	1	20050	825		WO 2	2005-1	EP15	73		2	0050	216
W: AE, AG,	AL, AM	, AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
CN, CO,	CR, CU	, CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
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AU 2005212830	A	1	20050	825		AU 2	2005-	2128	30		2	0050	216
CA 2555503													
EP 1716178													
R: AT, BE,													
	LT, LV		RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	HU,	PL,	SK,
BA, HR	IS, YU												
CN 1950399 BR 2005007649 ZA 2006006410	A		20070	1418		CN 2	2005-	B000	8594		2	0050	216
BR 2005007649	A		20070	710		BR 2	2005-	7649			2	0050	216
ZA 2006006410 JP 2008506353	A		20071	L227		ZA 2	2006-	5410			2	0050	216
JP 2008506353	Т		20080	306		JP 2	2006-	5525	76		2	0050	216
IN 2006CN02984	A		20070	0608		IN 2	2006-0	CN29	84		2	0060	814
IN 2006CN02984 MX 2006009253 NO 2006004183	A		20070	)418		MX 2	2006-	9253			2	0060	815
NO 2006004183	A		20061	1108		NO 2	006-	4183			2	0060	915
KR 2006131892	A		20061	1220		KR 2	006-	7189	76		2	0060	915
US 20080213256 PRAI EP 2004-3445	A	1	20080	)904		US 2	008-	5887.	34		2	0080	424
PRAI EP 2004-3445	A		20040	)216									
EP 2005-715354													
WO 2005-EP1573			20050			N T C	IIIO D	TODE		00143.5			

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
RE.ONI 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L4 ANSWER 8 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN
- TI Cloning and sequences of genetically engineered proteases that cleave target molecules for diagnosis or treatment of human diseases
- AB The present invention provides a disease treatment method by applying a medicament comprising a protease with defined target substrate specificity that enables hydrolysis of specific peptide bonds within the substrate related to such disease. This invention aims to create mutated proteases that target proteins or enzymes associated with disease (several dozen claimed mols.), for the purpose of hydrolysis-mediated alteration of cellular behavior aiding in diagnosis or treatment of human diseases. Specificity determining regions (SDR) from selected proteases were randomly inserted into a protein scaffold, enabling the protein scaffold to perform hydrolysis upon the SDR-determined substrate. Claimed are the sequences of human trypsin I, Bacillus subtiliss in E, human pepsin A, and human

caspase-7. Use of the modified trypsin protease upon tumor necrosis factor- $\alpha$ , serum proteins and VEGF, as well as anal. of corresponding cytotoxicity, is presented. The proteases with such a defined specificity can further be used for related therapeutic or diagnostic purposes.

2005:735080 HCAPLUS <<LOGINID::20100430>> AN

DN 143:206400

- ΤI Cloning and sequences of genetically engineered proteases that cleave target molecules for diagnosis or treatment of human diseases
- IN Haupts, Ulrich; Koltermann, Andre; Scheidig, Andreas; Votsmeier, Christian; Kettling, Ulrich; Coco, Wayne Michael

PA Germany

- SO U.S. Pat. Appl. Publ., 217 pp., Cont.-in-part of U.S. Ser. No. 872,198. CODEN: USXXCO
- DΤ Patent
- LA English

FAN.CNT 2

PAN.		TENT :	NO.			KIN		DATE			APPL	ICAT	ION	NO.		D	ATE	
PI		2005 1531 R:	179 AT,	BE,		A1 A1 DE,	DK,	2005 2005 ES, RO,	0518 FR, MK,	GB, CY,	EP 2 GR,		2587 LI,	LU,		SE,		111
		2004				A1		2004			AU 2	004-	2499	03		2	0040	618
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		2004		04		A1 A1		2004				004-					0040	
		2529				A1		2004				004-					0040	
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	EP	1633	865			A1		2006	0315		EP 2	004-	7418	41		2	0040	618
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	JP	2006			-,	T	,	2006			JP 2	006-	5161	70		2	0040	618
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PRAI	EP	2003	-138	19		A		2003	0618									

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EP 2003-25851 A 20031110
EP 2003-25871 A 20031111
US 2003-524960P P 20031125
US 2004-3058 A 20040211
US 2004-543518P P 20040211
US 2004-872198 A2 20040618
WO 2004-EP51172
                             W 20040618
WO 2004-EP51173
                             W
                                        20040618
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OSC.G THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD (3 CITINGS) 3

- ANSWER 9 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN
- ΤI Deimmunized fusion proteins comprising CD3-binding domain and Ig binding domain for diagnosis and treatment of cancer, inflammation, infection, (auto)immune disease or allergy
- AB The present invention provides a cytotoxically active CD3 specific binding construct comprising a first domain specifically binding to human CD3 and an Ig-derived second binding domain. Furthermore, a nucleic acid sequence encoding a CD3 specific binding construct of the invention is provided. The Iq.-derived binding domain comprises an antigen-interaction site with a specificity for mol. such as EpCAM, CCR5, CD19, Her-2, Her-2/neu, Her-3, Her-4, EGFR, PSMA, CEA, MUC-1, MUC2, MUC3, MUC4, MUC5AC, MUC5a, MUC7, BhCG, Lewis Y, CD20, CD33, CD30, GD3, 9-O-acetvl GD3, GM2, Globo H, fucosyl GM1, polySA, GD2, carboanhydrase IX, CD44v6, sonic Hedgehog, Wue-1, etc. Further aspects of the invention are vectors and host cells comprising said nucleic acid sequence, a process for the production of the construct of the invention and composition comprising said construct. The invention also provides the use of said constructs for the preparation of pharmaceutical compns. for the treatment of particular diseases, a method for the treatment of particular diseases and a kit comprising the binding construct of the invention.
- AN 2005:395357 HCAPLUS <<LOGINID::20100430>>
- DN 142:446010
- ΤI Deimmunized fusion proteins comprising CD3-binding domain and Ig binding domain for diagnosis and treatment of cancer, inflammation, infection, (auto)immune disease or allergy
- TN Hofmeister, Robert; Kohleisen, Birgit; Lenkkeri-Schuetz, Ulla; Itin, Christian; Baeuerle, Patrick; Carr, Francis J.; Hamilton, Anita A.; Williams, Stephen
- PA Micromet A.-G., Germany
  - PCT Int. Appl., 639 pp.
  - CODEN: PIXXD2
- DT Patent

LA FAN.		glish 1																
	PA:	ENT:	NO.			KIN	D	DATE			APPL	ICAT	ION :	NO.		D	ATE	
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								LV,										
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			SI,	SK,	TR,	BF,	ΒJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,
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	AU 2004283850					A1		2005	0506		AU 2	004-	2838	50		2	0041	015
	CA 2542239					A1		2005	0506		CA 2	004-	2542	239		2	0041	015
	EP 1673398					A1		2006	0628		EP 2	004-	7904	88		2	0041	015

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     CN 1867586
                               20061122 CN 2004-80030150
                         A
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                         С
                               20090121
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ZA 2006001699
JP 2007537714
                        A
                               20061205 BR 2004-15457
                                                                    20041015
                        A
T
                               20070530 ZA 2006-1699
                                                                    20041015
    JP 2007537714 T 20071227 JP 2006-534709 NZ 546173 A 20090130 NZ 2004-546173 NZ 506CN01280 A 20060831 MX 2006-001280 A 20070629 IN 2006-CN1280 NO 20060022117 A 20060703 NO 2006-22117
                                                                    20041015
                                                                    20041015
                                                                   20060410
                                                                   20060413
                                                                    20060511
US 20090022738 A1 20090122
PRAI EP 2003-23581 A 20031016
WO 2004-EP11646 W 20041015
                        A1 20090122 US 2006-572740
                                                                    20061204
ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OSC.G 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)
RE.CNT 6
             THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
     ANSWER 10 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN
TΙ
     Antibodies conjugated with phagocytic marker for enhancing phagocytosis
     against autoimmune disease, infection, cancer and others
AB
     The present invention provides a system for enhancing clearance or
     destruction of undesirable cells or noncellular mol. entities by tagging
     such cells or noncellular mol. entities with a marker that targets the
     cells or noncellular mol. entities for phagocytosis (phagocytic marker).
     The target cells can be, for example, endothelial cells, tumor cells,
     leukocytes, or virus-infected cells. In certain embodiments of the
     invention the tagging is accomplished by administering a composition comprising
     an antibody or ligand linked to the phagocytotic marker, wherein the
     antibody or ligand binds to a cell type specific marker present on or in
     the cell surface of a target cell. In preferred embodiments of the
     invention, the phagocytic marker comprises phosphatidylserine or a group
     derived from phosphatidylserine, thrombospondin-1, annexin I, or a derivative
     of any of these.
AN
    2005:182810 HCAPLUS <<LOGINID::20100430>>
DN
    142:278750
     Antibodies conjugated with phagocytic marker for enhancing phagocytosis
     against autoimmune disease, infection, cancer and others
TN
    Francois, Cedric; Olson, Paul; Deschatelets, Pascal; Machiels, Alec
    Potentia Pharmaceuticals, Inc., USA
PA
SO
    PCT Int. Appl., 173 pp.
     CODEN: PIXXD2
DT
     Patent.
LA
    English
FAN. CNT 1
     PATENT NO.
                        KIND DATE APPLICATION NO. DATE
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                                                                   -----
     WO 2005019429 A2 20050303 WO 2004-US27245
WO 2005019429 A3 20060302
PT
                                                                   20040823
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH,
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RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE,

SN. TD. TG

US	20050113297	A1	20050526	US	2004	1-923940		20040823	
PRAI US	2003-497086P	P	20030822						
US	2003-514941P	P	20031028						
US	2003-523611P	P	20031119						
US	2003-524126P	P	20031121						
US	2003-524730P	P	20031124						
US	2004-547951P	P	20040226						
WC	2004-US27245	A	20040823						
ASSIGNM	ENT HISTORY FOR	US PATENT	T AVAILABLE	IN I	LSUS	DISPLAY	FORMAT		
OS MA	RPAT 142:278750								

THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD (3 CITINGS) OSC.G 3 RE.CNT 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- T. 4 ANSWER 11 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN
- ΤТ Human molecules or antibodies specific to human CD3 complex for diagnosis and treatment of proliferative, inflammatory, immune, autoimmune, infectious or allergic diseases
- AB The present invention provides a method for the preparation of a human binding mol., fragment or derivative thereof which specifically binds to the human CD3 complex. The binding mols. are human, humanized or deimmunized antibodies or fragments; and are selected from a DNA or RNA library by a phage display method. The antibodies may comprise at least one further antigen-interaction-site and/or effector domain selected from EpCAM, CCR5, CD19, EphA2, HER-2 neu, HER-3, HER-4, EGFR, PSMA, CEA, MUC1, MUC2, MUC3, MUC4, MUC5, MUC7, BhCG, Lewis-Y, CD20, CD33, CD30, ganglioside GD3, These binding mols. or antibodies and fragments are useful for diagnosis and treatment of proliferative disease, tumor, inflammation, immune disease, autoimmune disease, infection, viral
  - infection, allergy, parasitic infection or graft vs. host disease.
- AN 2004:1059392 HCAPLUS <<LOGINID::20100430>>
- DN 142:36924
- ΤI Human molecules or antibodies specific to human CD3 complex for diagnosis and treatment of proliferative, inflammatory, immune, autoimmune, infectious or allergic diseases
- TN Kufer, Peter; Raum, Tobias; Berry, Meera; Kischel, Roman; Mangold, Susanne; Krinner, Eva; Kohleisen, Birgit; Zeman, Steven; Itin, Christian; Baeuerle, Patrick
- PA Micromet A.-G., Germany
- SO PCT Int. Appl., 350 pp. CODEN: PIXXD2
- DT Patent
- LA English

FAN.	CNT 1 PATENT	NO.		KIN	D	DATE			APPL	ICAT	ION I	NO.		D.	ATE	
PI	WO 2004			A2		2004			WO 2	004-1	EP56	84		2	0040	526
	WO 2004	106380	0	A3		2005	0623									
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		SN,	TD, TO													
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	CA 2523	716		A1		2004	1209		CA 2	004-	2523	716		2	0040	526

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EP 1629011 A2 20060301 EP 2004-739377 20040526
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    AT 455127
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                         20100115 AT 2004-739377
                                                        20040526
    IN 2005CN02915
                    A
                          20070914
                                    IN 2005-CN2915
                                                        20051108
    IN 228203
                    A1 20090306
PRAI EP 2003-12132
                    A 20030531
    WO 2004-EP5684
                    747
                          20040526
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THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD RE.CNT 1 ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L4 ANSWER 12 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN
- ΤI Glycoconjugates with NeuAc-NeuAc-Gal-Glc are more effective at preventing adhesion of Helicobacter pylori to gastric epithelial cells than glycoconjugates with NeuAc-Gal-Glc
- Helicobacter pylori (H. pylori) adheres to human gastric epithelial cells, AB eliciting various gastroduodenal diseases. Gangliosides play a critical role in bacterial adhesion to cell surfaces. The present study examined how residues of gangliosides are important for inhibition of adhesion of H. pylori to MKN-45 cells. We measured adhesion or detachment effects of cancliosides on the interaction between MKN-45 cells and H. pylori, as well as interleukin-8 production Among the gangliosides, O-Ac-GD3, GT1b, GD1a, GD1b, GT1a, and GD3 had potent dose dependent inhibitory effects on adhesion of H. pylori to MKN-45 cells, interleukin-8 production, and vacuole formation induced by H. pylori toxin binding to Vero cells. GD3 also accelerated bacterial detachment of MKN-45 cells with adherent H. pylori in a dose dependent manner. Such results strongly suggest that the mechanism involved in the inhibition of H. pylori adhesion is mediated by the variations of the residues of the NeuAc-NeuAc-Gal-Glc chain of gangliosides. NeuAc-NeuAc-Gal-Glc exhibits a more inhibitory effect on adhesion than the NeuAc-Gal-Glc chain. Such ganglioside and oligosaccharide sequences appear to have therapeutic importance for prevention of H. pylori adhesion, as well as reduction of both inflammation and gastric mucosal injuries.
- AN 2004:936243 HCAPLUS <<LOGINID::20100430>>
- DN 142:148329
- Glycoconjugates with NeuAc-NeuAc-Gal-Glc are more effective at preventing adhesion of Helicobacter pylori to gastric epithelial cells than glycoconjugates with NeuAc-Gal-Glc
- AU Hata, Y.; Murakami, M.; Okabe, S.
- Department of Geriatric Medicine, Kyoto University, Kyoto, Japan CS
- SO Journal of Physiology and Pharmacology (2004), 55(3), 607-625
- CODEN: JPHPEI: ISSN: 0867-5910
- PB Polish Physiological Society DT Journal
- T.A English
- OSC.G 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD (4 CITINGS) RE.CNT 58 THERE ARE 58 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- L4ANSWER 13 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN
- 36Fusion proteins comprising CD1d complex, a2 microglobulin and antibody or fragment for targeting therapy of tumor, autoimmune disease, inflammation and infection
- The invention is directed to a compound comprising one or more CDld complexes in association with an antibody specific for a cell surface marker. The CD1d complexes comprise a CD1d, a ss2-microglobulin mol., and may further comprise an antigen bound to the CD1d binding groove. The invention is further directed to methods of inhibiting or stimulating an immune response with the CDld-antibody compds., in particular anti-tumor

and autoimmunity responses.

- 2004:292071 HCAPLUS <<LOGINID::20100430>>
- DN 140:320040

AN

- 36Fusion proteins comprising CD1d complex, @2 microglobulin and antibody or fragment for targeting therapy of tumor, autoimmune disease, inflammation and infection
- IN Robert, Bruno; Donda, Alena; Cesson, Valerie; Mach, Jean-Pierre; Zauderer, Maurice
- PA Vaccinex, Inc., USA
- SO PCT Int. Appl., 152 pp.

WO 2003-US30238

IN 2005-KN523

- CODEN: PIXXD2
- Patent
- LA English

FAN.	CNT	1																
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PI		2004						2004			WO 2	003-	US30	238		2	0030	926
	WO	2004	0292	06		A3		2004	1007									
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	TN	2005																329
	US 20060269540						2006											
	IN 2007KN02053																	
PRAT	PRAI EP 2002-405838											007	20	-		-	0010	

20050329 OSC.G 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS) RE.CNT 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

20030926

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

ANSWER 14 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN L4

A3

- ΤI Antibodies produced by host cells tolerant to fucose-N-acetylglucosamine 1→6 α binding structure-recognizing lectins
- AB Disclosed is a process for producing an antibody composition with the use of cells tolerant to a lectin recognizing a sugar chain structure in which an α-bond is formed between the 6-position of N-acetylglucosamine at the reducing end of an N-glycoside bond-type complex sugar chain and the 1-position of fucose; and cells usable in this process. The antibodies exhibit enhanced antibody-dependent cytotoxicity. The host cells have lower or defective carbohydrate modification-related proteins such as (1) GDP-fucose synthesizing enzyme proteins, (2) fucose-N-acetylglucosamine 1→6 α-binding structure-modifying enzyme proteins, and (3) GDP-fucose to Golgi body-transporting proteins, e.g.

α-1,6-fucosyltransferase. The genes of these carbohydrate-modifying

enzymes are destroyed by gene targeting, dominant neg. body introduction, mutation or mutagenesis, transcription and/or translation inhibition, and RNAi. Antibodies prepared by the method include human antibodies, humanized or chimeric antibodies, antibody fragments and IgGs. These antibodies are prepared for diagnosis, prevention and treatment of cancer, allergy, inflammation, autoimmune disease, circulation disease, viral

infection and bacterial infection. AN 2003:818543 HCAPLUS <<LOGINID::20100430>>

DN 139:322290

- TI Antibodies produced by host cells tolerant to fucose-N-acetylglucosamine  $1\rightarrow6$   $\alpha$  binding structure-recognizing lectins
- Satoh, Mitsuo; Kamachi, Reiko; Kanda, Yutaka; Mori, Katsuhiro; Yamano, Kazuya; Kinoshita, Satoko; Iida, Shigeru
- PA Kyowa Hakko Kogyo Co., Ltd., Japan

SO PCT Int. Appl., 297 pp. CODEN: PIXXD2

DT Patent

LA Japanese FAN.CNT 1

1.4

	PA	TENT	NO.			KIN		DATE			APPL						ATE	
PI	WO	2003	0851	 18				2003:	1016								0030	409
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	AU	2003	2360	15		A1		2003	1020		AU 2	003-	2360	15		2	0030	409
	US	2004	0132	140		A1		2004	0708		US 2	003-	4096	16		2	0030	409
	EP	1498	490			A1		2005	0119		EP 2	003-	7230	96		2	0030	409
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
			IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL,	TR,	BG,	CZ,	EE,	HU,	SK	
PRAI	JP	2002	-106	820		A		2002	0409									
	JP	2003	-246	85		A		2003	0131									
	WO	2003	-JP4	502		W		2003	0409									
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OSC.G 17 THERE ARE 17 CAPLUS RECORDS THAT CITE THIS RECORD (35 CITINGS) RE.CNT 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- ANSWER 15 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN TI Antibodies produced by cells tolerant to lectin recognizing 1→6 α-bond between N-acetylglucosamine and fucose for diagnosis and therapy in patients suffering from FcyRIIIa polymorphism
- AΒ A drug containing, as the active ingredient, an antibody composition produced with

the use of cells tolerant to a lectin recognizing a sugar chain structure in which an α-bond is formed between the 6-position of N-acetylglucosamine at the reducing end of an N-glycoside bond-type complex sugar chain and the 1-position of fucose. This drug is appropriate for patients suffering from FcYRIIIa polymorphism who cannot be treated with a drug containing, as the active ingredient, an antibody composition produced from cells not tolerant to a lectin recognizing a sugar chain structure in which an α-bond is formed between the 6-position of N-acetylglucosamine at the reducing end of an N-glycoside

bond-type complex sugar chain and the 1-position of fucose. Such chimeric antibodies specific to GD3, FGF8, CD20, and CCR4 were prepared for diagnosis, prevention and treatment of tumor, allergy,

inflammation, autoimmune disease, circulation disorder, viral infection and bacterial infection.

AN 2003:818312 HCAPLUS <<LOGINID::20100430>>

DN 139:322285

- 7.1 Antibodies produced by cells tolerant to lectin recognizing 1-6 α-bond between N-acetylglucosamine and fucose for diagnosis and therapy in patients suffering from FcγRIIIa polymorphism IN Nakamura, Kazuyasu; Shitara, Kenya; Hatanaka, Shigeki; Niwa, Rinpei;
- IN Nakamura, Kazuyasu; Shitara, Kenya; Hatanaka, Shigeki; Niwa, Rinpei; Okazaki, Akira
- PA Kyowa Hakko Kogyo Co., Ltd., Japan

SO PCT Int. Appl., 214 pp. CODEN: PIXXD2

DT Patent

LA Japanese

FAN CNT 1

FAN.	PATENT	NO.			KIN	D	DATE			APPL	ICAT	ION	NO.		D	ATE	
PI	WO 200	30845	70		A1		2003	1016		WO 2	003-	JP45	05		2	0030	409
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							ΙE,										
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	CA 248						2003										
	AU 200						2003										
	EP 150		2603														
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	US 20050031613				A1		2005			US 2	003-	4096	08		2	0030	109
PRAI	AI JP 2002-106951						2002										
	WO 2003-JP4505						2003	0409									

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
RE.CNT 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L4 ANSWER 16 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN
- TI Isolation and identification of buffalo milk gangliosides and their use for humanization of infant and other formulas
- AB The present invention relates to gangliosides derived or isolated from buffalo milk, skimmed buffalo milk, buffalo milk serum or derivs. of either. Buffalo milk is reported to comprise gangliosides that are not contained in bovine milk, such as gangliosides that belong to the GMI-class. Furthermore, buffalo milk is found to comprise unknown gangliosides, denoted herein as ganglioside "F" and "L". Furthermore, the invention reports that gangliosides are surprisingly found in fractions of isolation procedures that were so far not considered to comprise gangliosides. Finally, milk or milk serum from buffalo, for example as derived from mozzarella cheese production, contains specific gangliosides in the same amts. as human breast milk, which makes it suitable for humanization of infant and other formulas. Anti-inflammatory effects of buffalo milk gangliosides are also disclosed.

AN 2003:509876 HCAPLUS <<LOGINID::20100430>>

DN 139:68312

- TI Isolation and identification of buffalo milk gangliosides and their use for humanization of infant and other formulas
- IN Colarow, Ladislas; Turini, Marco; Berger, Alvin
- PA Societe des Produits Nestle S.A., Switz.
- SO Eur. Pat. Appl., 24 pp. CODEN: EPXXDW
  - T Patent
- LA English

FAN.CNT 1

PAN.																		
		TENT						DATE									ATE	
PI		1323															0011	227
PI	EP																	
		R:						ES,					LI,	LU,	NL,	SE,	MC,	PT,
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	WO	2003	0554	97		A1		2003	0710		WO 2	002-	EP14	876		2	0021	220
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PRAI	EP	2001	-130	614		A		2001	1227									
		2002																
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OSC.G 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)
RE.CNT 14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L4 ANSWER 17 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN
- TI Inducing tolerance or immunomodulation using dendritic cells incubated with antigen
  - Disclosed is a method of altering immune responses using dendritic cells. One form of the method is a method of inducing immunol, tolerance in an individual, where type 2 dendritic cells are administered to an individual, and where the dendritic cells have been incubated with one or more antigens. Another form of the method involves altering an immune response, in which liposomes containing one or more antigens are administered to an individual, and where the liposomes are modified with surface-bound mols. that target the liposomes to type 2 dendritic cells. Another form of the method involves reducing immune responsiveness, where liposomes containing one or more antigens are administered to an individual and where the liposomes are modified with the surface bound mols, that target the liposomes to type 1 dendritic cells or type 2 dendritic cells. Another form of the method is a method of enhancing immune responsiveness, where liposomes containing one or more antigens are administered to an individual, and where the liposomes are modified with surface-bound mols. that target the liposomes to mature type 1 dendritic cells. The antigens can be autoantigens, alloantigens, tumor antigens, and viral antigens, and can be in the form of carbohydrates, peptides, nucleic acids, and lipids. The liposome surface-bound mols. can be specific for CD11c+ and/or BDCA-1,

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which targets mature type 1 dendritic cells. Type 2 dendritic cells can
     be targeted by using surface-bound mols. specific for CD123, BDCA-2,
     and/or BDCA-4.
     2002:869052 HCAPLUS <<LOGINID::20100430>>
    137:336727
    Inducing tolerance or immunomodulation using dendritic cells incubated
     with antigen
    Waller, Edmund K.; Rosenthal, Hillary S.; Lonail, Sagar
    Emory University, USA
     PCT Int. Appl., 34 pp.
     CODEN: PIXXD2
     Patent
     English
FAN.CNT 1
     PATENT NO.
                       KIND DATE
                                          APPLICATION NO.
                                                                  DATE
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    WO 2002090510
                        A2
                              20021114 WO 2002-US14497
                                                                  20020508
                        A3
     WO 2002090510
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     WO 2002090510
                        A9 20040429
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     AU 2002305452
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                              20021118 AU 2002-305452
                                                                   20020508
     US 20050013810
                         A1
                               20050120
                                           US 2004-477012
                                                                   20040430
PRAI US 2001-289625P
                         P
                                20010508
    WO 2002-US14497
                         W
                                20020508
OSC.G
             THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)
    ANSWER 18 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN
    Colostrum-based pharmaceutical compositions
    A composition including colostrum or a colostrum-derived product and
     hyperimmune milk (HIM) or a hyperimmune milk-derived product, in amts.
     sufficient to provide a combined spectrum of pathogen-binding activity
     against a broad-spectrum of pathogenic organisms is described. For
     example, a test composition was prepared including 70% colostrum milk protein
     powder, 24% hyperimmune milk powder, 4% ganglioside-containing component, whey
     powder, lactose and 1.5% milk calcium. The test composition of the invention
     includes a combination of ingredients each of which has particular
     antimicrobial binding and/or anti-inflammatory activity which
    may combine to produce particular and unexpected clin. benefits in a broad
    range of diseases, including infection-associated diseases, and particularly
     gastrointestinal, inflammatory and bone related disorders. Such
     benefits are an unexpected result of the combination used.
     2002:391563 HCAPLUS <<LOGINID::20100430>>
     136:391021
    Colostrum-based pharmaceutical compositions
    Williams, Charles Edward; Hobman, Peter Graeme; Yarrow, Simon Stephen
    Fonterra Co-Operative Group Limited, N. Z.
    PCT Int. Appl., 43 pp.
    CODEN: PIXXD2
    Patent
    English
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FAN.CNT 1 PATENT NO. KIND DATE APPLICATION NO. DATE

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A1 20020523 WO 2001-NZ256 20011115
    WO 2002040051
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20030910 EP 2001-996393 20011115
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    EP 1341554
                         A1
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    HU 2004000589
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                                                                 20011115
    HU 2004000589
                        A3 20050628
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    CN 1299771
                              20070214
                                          CN 2001-822044
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    US 20040047856
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A1
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20051006
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PRAI NZ 2000-508234
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            THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD (2 CITINGS)
OSC.G
RE.CNT 3
             THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD
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ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 19 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN TI Novel synthetic gangliosides

TI

AB Disclosed are novel synthetic ganglioside comprising a modified sphingosine group represented by Structural Formula (I); Y is -O- or -NH-; X is =0 or -H2; R1 and R2 are independently a substituted or unsubstituted straight chain or branched hydrocarbyl group, wherein the hydrocarbyl group optionally comprises -S-, -S(0)-, -S02-, -O- or -NR- (each R is independently -H, an aliphatic group, a substituted aliphatic group, an aryl group or a substituted aryl group); and R3 is -H, -S(O)2H, -P(O)2OH, -N(O)OH or -P(O)2OP(O2)OH. Also disclosed are methods of treating a subject with a neurol. condition or disease and methods of treating a subject in need of immunosuppression. The subject can be, e.g., in need of neuroprotection, in need of neurogenesis, or in need of neuritogenesis. The method can be used for immunosuppression, e.g., a subject with organ, bone marrow, or stem cell transplant or a subject with autoimmune disease. The methods comprises the step of administering to the subject an effective amount of the synthetic ganglioside represented by Structural Formula (I).

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AN
     2002:171915 HCAPLUS <<LOGINID::20100430>>
DN
     136:210593
TT
     Novel synthetic gangliosides
IN
    Ho, Tony W.
PA
    Neuronyx, Inc., USA
     PCT Int. Appl., 38 pp.
SO
     CODEN: PIXXD2
     Patent
LA
     English
FAN.CNT 1
                        KIND DATE APPLICATION NO.
ΡI
     WO 2002018401
                         A2 20020307 WO 2001-US27087
A3 20020822
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     AU 2001085359
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PRAI US 2000-654363
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     WO 2001-US27087
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    MARPAT 136:210593
OSC.G 1 THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD (1 CITINGS)
RE.CNT 1
              THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD
              ALL CITATIONS AVAILABLE IN THE RE FORMAT
     ANSWER 20 OF 20 HCAPLUS COPYRIGHT 2010 ACS on STN
L4
     Methods for treatment of tumors and metastases using a combination of
ΤI
     anti-angiogenic and immunotherapies
AB
     The invention teaches methods for treating tumors and tumor metastases in
     a mammal comprising administering, to a mammal in need of treatment, a
     therapeutic amount of an antagonist sufficient to inhibit angiogenesis in
     combination with a therapeutic amount of anti-tumor immunotherapeutic agent,
     such as an anti-tumor antigen antibody/cytokine fusion protein having a
     cytokine and a recombinant Iq polypeptide chain sufficient to elicit a
     cvtokine-specific biol. response.
AN
    2000:573686 HCAPLUS <<LOGINID::20100430>>
DN
    133:176175
     Methods for treatment of tumors and metastases using a combination of
     anti-angiogenic and immunotherapies
     Lode, Holger N.; Reisfeld, Ralph A.; Cheresh, David A.; Gillies, Stephen
TN
PA
     The Scripps Research Institute, USA; Lexigen Pharmaceuticals Corporation
SO
     PCT Int. Appl., 78 pp.
     CODEN: PIXXD2
     Patent
LA
    English
FAN.CNT 1
                        KIND DATE APPLICATION NO. DATE
     PATENT NO.
     WO 2000047228 A1 20000817 WO 2000-US3483 20000211
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                HU 2002000128 A2 20020529 HU 2002-128 PU 2002-128 PU 2002536419 T 20021029 JP 2000-598179 RU 2236251 C2 20040920 RU 2001-124907 CN 1192796 C 20050316 CN 2000-806134 PU 3715261 B1 20061003 US 2000-502732 AT 412433 T 20081115 AT 2009-910138 PT 1156823 E 20090108 PT 2000-910138 ES 2313883 T3 20090316 ES 2000-910138 ES 2313883 T3 20090316 ES 2000-910138 CN 2001006455 A 2002106 ZA 2001-6455 PU 2009-9108 PU 200
                  HU 2002000128
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OSC.G 4 THERE ARE 4 CAPLUS RECORDS THAT CITE THIS RECORD (5 CITINGS)
RE.CNT 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> s cholesterol or hypercholesterolem? or atherosclerosis 211313 CHOLESTEROL

20859 HYPERCHOLESTEROLEM?

72634 ATHEROSCLEROSIS

265180 CHOLESTEROL OR HYPERCHOLESTEROLEM? OR ATHEROSCLEROSIS

=> s 12 and 15 L6 8 L2 AND L5

1.5

=> d 16 1-8 ti abs bib

L6 ANSWER 1 OF 8 HCAPLUS COPYRIGHT 2010 ACS on STN

- TI Method and apparatus of low strength electric field network-mediated delivery of drug, gene, siRNA, shRNA, protein, peptide, antibody or other biomedical and therapeutic molecules and reagents in skin, soft tissue, joints and bone
- AB The invention includes four preferred embodiments: (i) a method and apparatus for the joint and its related soft tissue for bone gene, protein and drug delivery; (ii) a method and apparatus for gene, protein and drug delivery to its method and apparatus for delivery of gene, protein and drug delivery to skin and soft tissue; and/or (iv) a method and apparatus for delivery of a gene, protein and drug to soft tissue tumor. The apparatus for transfecting drug, gene, siRNA, shRNA, protein, peptide, antibody or other biomedical and therapeutic mols. and reagents comprises a plurality of neg. electrodes disposed into low resistance elec. contact with skin overlaying the tissue.

- AN 2007:1207931 HCAPLUS <<LOGINID::20100430>>
- DN 147:474740
- ΤТ Method and apparatus of low strength electric field network-mediated delivery of drug, gene, siRNA, shRNA, protein, peptide, antibody or other biomedical and therapeutic molecules and reagents in skin, soft tissue, joints and bone
- ΤN Sen, Luyi
- PA The University of California, USA
- SO PCT Int. Appl., 51 pp.
- CODEN: PIXXD2 DT Patent
- LA English

	N.CNT 1																		
	PA:	ENT I	NO.			KIN		DATE			APPL	ICAT	ION I	NO.		D	ATE		
		2007				A2			1025 1113		WO 2	007-	JS84	45		2	0070	402	
		W:	CH,	CN,	co,	AM, CR, GM,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	
			KN, MK,	ΚΡ, MN,	KR, MW,	KZ, MX, SC,	LA, MY,	LC, MZ,	LK, NA,	LR, NG,	LS, NI,	LT, NO,	LU, NZ,	LY, OM,	MA, PG,	MD, PH,	ME, PL,	MG, PT,	
		RW:	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	ZW					IE,	
			ВJ,	CF,	CG,	LU,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,	
			BY,	KG,	KZ,	LS, MD,	RU,	TJ,	TM,	AP,	EA,	EP,	OA						
		2647				A1 A2													
			AT, IS,	BE, IT,	BG, LI,	CH, LT, MK,	CY, LU,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,	
	IN	1015 2008	06370 CN05	) 122	·	A A		2009	0812 0320								0070 0081		
	US	2006- 2006- 2007-	-8192	277P		P		2006	0706										
OSC.G	3	1	THE	ERE A	ARE	1 CA	PLUS	REC	ORDS	THA	T CI	TE T	HIS I	RECO	RD (	1 CI	TING:	S)	

- L6 ANSWER 2 OF 8 HCAPLUS COPYRIGHT 2010 ACS on STN
- TΙ Formulations for mediating inflammatory bowel disorders
- The invention provides formulations and methods for mediating inflammation, in particular an inflammatory bowel disorder such as necrotizing enterocolitis. Further, the formulations are effective in lowering blood cholesterol and decreasing blood cholesterol absorption. The formulations comprise at least one ganglioside, which may be selected from the group consisting of: GD3, GM1, GM2, GM3, and GD1b. The invention provides a method of treating or preventing inflammatory diseases, such as necrotizing enterocolitis by delivery of at least one ganglioside to a subject in need thereof. Supplementation of foods or ligs, with gangliosides, for example infant formula or infant foods, can be employed according to the invention.
- 2007:815148 HCAPLUS <<LOGINID::20100430>> AN
- DN 147:197354
- TI Formulations for mediating inflammatory bowel disorders
- TN Clandinin, Michael Thomas; Park, Eek J.
- PΑ Mti Meta Tech Inc., Can.
- SO U.S. Pat. Appl. Publ., 39pp., Cont.-in-part of U.S. Ser. No. 551,789 CODEN: USXXCO

DT Patent LA English FAN.CNT 2

	PA:	TENT NO.				KIND DATE			APPLICATION NO.						DATE			
PI	WO				A2		2007 2004 2004	1014										
		W:	AE, CN, GE, LK, NO, TJ, BW, BY,	AG, CO, GH, LR, NZ, TM, GH, KG, FI, TR,	AL, CR, GM, LS, OM, TN, GM, KZ, FR,	AM, CU, HR, LT, PG, TR, KE, MD, GB,	AT, CZ, HU, LU, PH, TT, LS, RU, GR,	AU, DE, ID, LV, PL, TZ, MW, TJ, HU, CG,	AZ, DK, IL, MA, PT, UA, MZ, TM, IE,	DM, IN, MD, RO, UG, SD, AT, IT,	DZ, IS, MG, RU, US, SL, BE, LU,	EC, JP, MK, SC, UZ, SZ, BG, MC,	EE, KE, MN, SD, VC, TZ, CH, NL,	EG, KG, MW, SE, VN, UG, CY, PL,	ES, KP, MX, SG, YU, ZM, CZ, PT,	FI, KR, MZ, SK, ZA, ZW, DE, RO,	GB, KZ, NA, SL, ZM, AM, DK, SE,	GD, LC, NI, SY, ZW AZ, EE, SI,
PRAI	US WO	2006 2004 2004 2003	0276 -551 -CA3	430 789 75		W		2006 2004 2004 2003	0312 0312		US 2	004-	5517	89		2	0040	312

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

L6 ANSWER 3 OF 8 HCAPLUS COPYRIGHT 2010 ACS on STN

TI Anti-atherosclerotic mechanisms of ganglioside GD3, antioxidant PDTC and flavonoid quercetin in vascular smooth muscle cells

AB A review. Sialic acid containing glycosphingolipids (gangliosides) have been implicated in regulating various biol. phenomena such as atherosclerosis. Disialoganglioside (GD3) inhibited DNA synthesis of cultured VSMC in the presence of PDGF with down-regulation of cyclinE/CDK2 and up-regulation of the CDK inhibitor p21 and p27 expression. GD3 inhibited TNF-α-induced matrix metalloproteinase-9 (MMP-9) expression in VSMC and decreased MMP-9 promoter activity in response to TNF- $\alpha$ , which was transcriptionally regulated at NF-KB and activation protein-1 (AP-1) sites in the MMP-9 promoter. These suggest that the GD3 represents a physiol. modulator of VSMC responses that may contribute to plaque instability in atherosclerosis. On the other hand, pyrrolidine dithiocarbamate (PDTC), a metal chelating antioxidant and pro-oxidant compound reduced cell growth and DNA synthesis on VSMC in low d. conditions. However, in serum depleted medium, PDTC did not affect the cell viability. At low VSMC d. in 10% FBS, PDTC induced cell cycle arrest in the G1 phase. The cell cycle arrest is associated with the down-regulation of cyclin D1, cyclin E, CDK2, CDK4 and up-regulation of the CDK inhibitor p21 expression. These inhibitory effects were associated with enhanced expression of p21 and increased complexing of p21 with cyclin D1/CDK4 and cyclin E/CDK2. PDTC induced marked activation of p38MAPK and JNK. SB203580, a p38MAPK specific inhibitor, blocked PDTC-dependent p38MAPK, growth inhibition, and p21 expression. The cells were transfected with antisense-p21 oligodeoxynucleotide also decreased PDTC-induced p38 MAPK activity. These data demonstrate that the p38MAPK pathway participates in p21 induction, leading to decrease of cyclin D1/cdk4 and cyclin E/cdk2 complexes and PDTC-dependent VSMC growth inhibition. Finally, quercetin, a bioflavonoid, is known to inhibit angiotensin II-induced hypertrophy and serum-induced smooth muscle cell proliferation. Treatment of quercetin showed potent inhibitory effects on DNA synthesis of cultured human aortic smooth muscle cells (HASMC) in the presence of TNF- $\alpha$ . These inhibitory effects were associated with reduced extracellular signal-regulated kinase (ERK) 1/2 activity and G1 cell cycle arrest.

Quercetin induced down-regulation of cyclins and CDKs and up-regulation of the CDK inhibitor p21 expression. Quercetin inhibited TNF- $\alpha$ -induced MMP-9 secretion on HASMC in a dose dependent manner by down-regulation of MMP-9, indicating the efficacy of quercetin in inhibiting cell proliferation, G1 to S phase cell cycle progress and MMP-9 expression through the transcription factors NF-kB and AP-1 on TNF-α-induced HASMC.

- AΝ 2007:413849 HCAPLUS <<LOGINID::20100430>>
- DN 146:513511
- TI Anti-atherosclerotic mechanisms of cancilioside GD3, antioxidant PDTC and flavonoid quercetin in vascular smooth muscle cells
- Kim, Cheorl-Ho; Jin, Un-Ho; Suh, Seok-Jong; Moon, Sung-Kwon
- CS National Research Laboratory for Glycobiology, Korean Ministry of Science and Technology, Kyungju, 780-714, S. Korea SO Current Topics in Biotechnology (2005), 2, 93-113
- CODEN: CTBUAI; ISSN: 0972-821X
- Research Trends PR
- DT Journal; General Review
- LA English
- RE.CNT 77 THERE ARE 77 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT
- ANSWER 4 OF 8 HCAPLUS COPYRIGHT 2010 ACS on STN
- ΤI Methods of cancer treatment/prevention using cancer cell-specific surface antigens
- AB The authors disclose the elicitation of specific cellular and humoral immune responses against cancer cell surface antigens, including those cancer cell surface antigens expressed only in cancer cells and in non-cancer cells normally located in one or more immune-privileged sites or tissues of the individual. The method comprises using specifically prepared immunogen in fresh or lyophilized liposomes, proper routes of administration of the immunogen, proper doses of the immunogen, and specific combinations of heterologous immunization including DNA priming followed by liposomal protein boost to tailor the immune responses. In one example, the authors employ liposomal HBsAg as a model cancer antigen. AN
  - 2006:438034 HCAPLUS <<LOGINID::20100430>>
- DN 144:449376
- Methods of cancer treatment/prevention using cancer cell-specific surface
- Kislauskis, Edward; Yang, Kejian; Whalen, Barbara J.
- Biomedical Research Models, Inc., USA PA
- SO PCT Int. Appl., 85 pp. CODEN: PIXXD2
- DT Patent. LA English
- FAN. CNT 1

	PATENT NO.					KIN	D	DATE			APPLICATION NO.						DATE		
							_									-			
PI	WO	2006	0501	16		A1		2006	0511	1	WO 2	005-	US38	968		2	0051	027	
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			CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
			GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,	KΡ,	KR,	
			KZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	
			MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	
			SG,	SK,	SL,	SM,	SY,	ΤJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	
			VN,	YU,	ZA,	ZM,	zw												
		RW.	AT.	BE.	BG.	CH.	CY.	CZ.	DE.	DK.	EE.	ES.	FT.	FR.	GB.	GR.	HII.	TE.	

RW: AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG, BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

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EP 1812052 A1 20070801 EP 2005-819700 20051027
R: AT, BE, BG, CH, CY, CZ, DE, DK, EB, ES, FI, FR, GB, GR, HU, IE,
IS, IT, LI, LT, LU, LV, MC, NL, PL, PT, RO, SE, SI, SK, TR
US 20080267963 A1 20081030 US 2008-666956 20080521
PRAI US 2004-624296P P 20041102
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WO 2005-US38968 W 20051027

RE.CNT 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 5 OF 8 HCAPLUS COPYRIGHT 2010 ACS on STN

TI Antibodies conjugated with phagocytic marker for enhancing phagocytosis against autoimmune disease, infection, cancer and others

- AB The present invention provides a system for enhancing clearance or destruction of undesirable cells or noncellular mol. entities by tagging such cells or noncellular mol. entities with a marker that targets the cells or noncellular mol. entities for phagocytosis (phagocytic marker). The target cells can be, for example, endothelial cells, tumor cells, leukocytes, or virus-infected cells. In certain embodiments of the invention the tagging is accomplished by administering a composition comprising an antibody or ligand linked to the phagocytotic marker, wherein the antibody or ligand binds to a cell type specific marker present on or in the cell surface of a target cell. In preferred embodiments of the invention, the phagocytic marker comprises phosphatidylserine or a group derived from phosphatidylserine, thrombospondin-1, annexin I, or a derivative of any of these.
- AN 2005:182810 HCAPLUS <<LOGINID::20100430>>

DN 142:278750

- I Antibodies conjugated with phagocytic marker for enhancing phagocytosis against autoimmune disease, infection, cancer and others
- IN Francois, Cedric; Olson, Paul; Deschatelets, Pascal; Machiels, Alec
- PA Potentia Pharmaceuticals, Inc., USA
- SO PCT Int. Appl., 173 pp. CODEN: PIXXD2
- DT Patent
- LA English

		ENT I				KIN	D			APPLICATION NO.								
ΡI	WO	2005	0194	29				2005	0303								0040	
		W: RW:	CN, GE, LK, NO, TJ, BW, AZ, EE, SI,	CO, GH, LR, NZ, TM, GH, BY, ES, SK,	CR, GM, LS, OM, TN, GM, KG, FI, TR,	CU, HR, LT, PG, TR, KE, KZ,	CZ, HU, LU, PH, TT, LS, MD, GB,	AU, DE, ID, LV, PL, TZ, MW, RU, GR, CF,	DK, IL, MA, PT, UA, MZ, TJ, HU,	DM, IN, MD, RO, UG, NA, TM, IE,	DZ, IS, MG, RU, US, SD, AT, IT,	EC, JP, MK, SC, UZ, SL, BE, LU,	EE, KE, MN, SD, VC, SZ, BG, MC,	EG, KG, MW, SE, VN, TZ, CH, NL,	ES, KP, MX, SG, YU, UG, CY, PL,	FI, KR, MZ, SK, ZA, ZM, CZ, PT,	GB, KZ, NA, SL, ZM, ZW, DE, RO,	GD, LC, NI, SY, ZW AM, DK, SE,
PRAI	US US US US US	2005 2003 2003 2003 2003 2003	0113: -497: -514: -523: -524: -524:	086P 941P 611P 126P 730P		P P P P		2005 2003 2003 2003 2003 2003 2003	0822 1028 1119 1121	1	US 2	004-	9239	40		2	0040	823
		2004				P A		2004 2004										

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

OSC.G 3 THERE ARE 3 CAPLUS RECORDS THAT CITE THIS RECORD (3 CITINGS) RE.CNT 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 6 OF 8 HCAPLUS COPYRIGHT 2010 ACS on STN

TI Potentiation of immune responses with liposomal adjuvants

AB A high-integrity liposome comprising at least one stable lipid and at least one peptide-like therapeutic agent associated with said liposome, adapted for parenteral administration to an animal, including a human, and method according to manufacture and use are disclosed. Immunizing dosage forms comprising a liposome and an immunogen, wherein said liposome and immunogen are present in an immunization dose are provided. Addnl., a dosage form, including such form particularly adapted to producing an immune response, comprising a salt according to an organic acid derivative of a sterol and an immunogen wherein said organic acid derivative of a sterol and immunogen are present in an immunization dose, and method according to use is disclosed. Further, a dosage form, including such form particularly adapted to producing an immune response, comprising dimyristoylphosphatidylcholine (DMPC)/cholesterol liposomes, optionally in an aluminum hydroxide gel, and an immunogen wherein said DMPC/cholesterol and immunogen are present in an immunization dose, and method according to use is presented.

AN 2000:492029 HCAPLUS <<LOGINID::20100430>>

DN 133:109954

Potentiation of immune responses with liposomal adjuvants

Popescu, Mircea C.; Weiner, Alan L.; Recine, Marie S.; Janoff, Andrew S.; Estis, Leonard; Keyes, Lynn D.; Alving, Carl R.

PA The Liposome Company, Inc., USA

SO U.S., 23 pp., Cont.-in-part of U.S. 5,231,112.

CODEN: USXXAM

DТ Patent English

LA EAN ONT

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 6090406	Α	20000718	US 1990-485388	19900226
	US 4721612	A	19880126	US 1985-721630	19850410
	JP 09040550	A	19970210	JP 1996-191707	19850411
	US 4891208	A	19900102	US 1985-773429	19850910
	ZA 8507576	A	19860625	ZA 1985-7576	19851001
	IL 96444	A	19921201	IL 1985-96444	19851006
	DD 255533	A5	19880406	DD 1985-281616	19851010
	AU 8775438	A	19880111	AU 1987-75438	19870612
	JP 01501622	T	19890608	JP 1987-503771	19870612
	CA 1337898	С	19960109	CA 1988-584808	19881202
	US 6759057	B1	20040706	US 1989-323182	19890313
	AU 8941861	A	19900323	AU 1989-41861	19890824
	AU 627226	B2	19920820		
	AU 8942214	A	19900323	AU 1989-42214	19890824
	AU 631377	B2	19921126		
	JP 04500203	T	19920116	JP 1989-509162	19890824
	CA 1334165	С	19950131	CA 1989-609463	19890825
	US 5231112	A	19930727	US 1989-425727	19891023
	JP 07100367	A	19950418	JP 1993-268664	19931027
	JP 2568034	B2	19961225		
	US 5897873	A	19990427	US 1995-392676	19950223
PRA	I US 1984-599691	B2	19840412		
	US 1985-721630	A2	19850410		
	US 1985-773429	A2	19850910		
	US 1986-873584	B2	19860612		
	US 1986-934151	B2	19861124		

US	1987-61186	B2	19870611
US	1987-128974	B2	19871204
US	1988-236701	B2	19880825
US	1988-236702	B2	19880825
US	1988-277854	B2	19881130
US	1989-397777	B2	19890823
US	1989-425727	A2	19891023
JP	1985-502090		19850411
JP	1993-268664	A3	19850411
IL	1985-76600	A3	19851006
WO	1987-US1402	A	19870612
US	1989-397758	A	19890823
WO	1989-US3657	A	19890824
WO	1989-US3658	A	19890824
US	1991-758587	A1	19910912
US	1993-108822	A2	19930818
US	1993-146463	B1	19931102

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT OSC.G 41 THERE ARE 41 CAPLUS RECORDS THAT CITE THIS RECORD (47 CITINGS) RE.CNT 62 THERE ARE 62 CITED REFERENCES AVAILABLE FOR THIS RECORD ALL CITATIONS AVAILABLE IN THE RE FORMAT

- ANSWER 7 OF 8 HCAPLUS COPYRIGHT 2010 ACS on STN L6
- ΤI Peptide-containing liposomes, immunogenic liposomes and methods of preparation and use
- AB A high integrity liposome comprising at least one stable lipid and at least one peptide-like therapeutic agent associated with the liposome, adapted for parenteral administration to an animal, including a human, and a method for manufacture and use are disclosed. Immunizing dosage forms comprise a liposome and an immunogen, wherein the liposome and immunogen are present in an immunization dose. Addnl., a dosage form, including such form particularly adapted to producing an immune response, comprising a salt according to an organic acid derivative of a sterol and an immunogen present in an immunization dose, and a method for use are disclosed. Further, a dosage form, including such form particularly adapted to producing an immune response, comprising dimyristolyphosphatidylcholine (DMPC)/cholesterol liposomes, optionally in an aluminum hydroxide gel, and an immunogen wherein the DMPC/cholesterol and immunogen are present in an immunization dose, and method for their use are disclosed.
- 1999:412601 HCAPLUS <<LOGINID::20100430>> AN
- DN 131:63430
- ΤI Peptide-containing liposomes, immunogenic liposomes and methods of preparation and use
- IN Popescu, Mircea C.; Weiner, Alan L.; Recine, Marie S.; Janoff, Andrew S.; Estis, Leonard; Keyes, Lynn D.; Alving, Carl R. PA
- The Liposome Company, Inc., USA
- SO U.S., 24 pp., Cont.-in-part of U.S. Ser. No. 108,822. CODEN: USXXAM
- DT Patent.
- LA English FAN.CNT 9

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
PI	US 5916588	A	19990629	US 1995-452549	19950525
	US 4721612	A	19880126	US 1985-721630	19850410
	JP 09040550	A	19970210	JP 1996-191707	19850411
	US 4891208	A	19900102	US 1985-773429	19850910
	ZA 8507576	A	19860625	ZA 1985-7576	19851001
	IL 96444	A	19921201	IL 1985-96444	19851006
	DD 255533	A5	19880406	DD 1985-281616	19851010

	2.11	8775438		-	19880111	2.55	1987-75438		19870612
				A					
		01501622		T C	19890608		1987-503771 1988-584808		19870612
		1337898			19960109				19881202
		6759057		B1	20040706		1989-323182		19890313
		8941861		A	19900323	AU	1989-41861		19890824
		627226		B2	19920820		4000 40044		
		8942214		A	19900323	AU	1989-42214		19890824
		631377		B2	19921126				
		04500203		T	19920116		1989-509162		19890824
		1334165		C	19950131		1989-609463		19890825
		5231112		A	19930727		1989-425727		19891023
		5288499		A	19940222		1991-758587		19910912
		6352716		B1	20020305		1993-108822		19930818
		07100367		A	19950418	JP	1993-268664		19931027
		2568034		B2	19961225				
		5897873		A	19990427	US	1995-392676		19950223
PRAI		1984-599691		B2	19840412				
		1985-721630		A2	19850410				
		1985-773429		A2	19850910				
	US	1986-873584		B2	19860612				
	US	1986-934151		A2	19861124				
		1987-61186		B2	19870611				
		1987-128974		B2	19871204				
		1988-236701		A2	19880825				
		1988-236702		B2	19880825				
		1988-277854		B2	19881130				
	US	1989-397777		B2	19890823				
	US	1989-425727		A3	19891023				
		1991-758587		A1	19910912				
		1993-108822		A2	19930818				
	JP	1985-502090			19850411				
	JP	1993-268664		A3	19850411				
	IL	1985-76600		A3	19851006				
	WO	1987-US1402		A	19870612				
		1989-397758		A	19890823				
	WO	1989-US3657		A	19890824				
	WO	1989-US3658		A	19890824				
	US	1993-146463		B1	19931102				
A C C T C	TAILE	DOM VECTORY FOR	TTC	DATES	OTOSTESUS TE	TAT	CITC DICDIAN	EODMAT.	

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OSC.G 5 THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD (7 CITINGS)
RE.CNT 60 THERE ARE 60 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

- L6 ANSWER 8 OF 8 HCAPLUS COPYRIGHT 2010 ACS on STN
- TI Ganglioside immunostimulating complexes and uses thereof
- AB The present invention relates generally to an immunostimulating complex comprising one or more gangliosides and more particularly to an immunostimulating complex comprising at least one of the gangliosides GM2, GD2, GD3 or GT3. The immunogenic immunostimulating complex also comprises a saponin preparation, a sterol, a protein epitope, and phospholipid. The protein may be cancer specific protein, melanoma specific protein, or influenza hemagquitainn. The present invention is useful, inter alla, as a prophylactic and/or therapeutic agent in the treatment of tumors, and more particularly, melanomas.
- AN 1999:7859 HCAPLUS <<LOGINID::20100430>>
- DN 130:65237
- TI Ganglioside immunostimulating complexes and uses thereof
- IN Cox, John Cooper; Ronnberg, Bengt John Lennart; Sjolander, Sigrid Elisabet
- PA Eriksson, Lennart, Australia; CSL Limited
- SO PCT Int. Appl., 40 pp.

CODEN: PIXXD2

FAN.	FAN.CNT 1 PATENT NO.										APPLICATION NO.								
PI												1998-							
		W:	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR	, BY,	CA,	CH,	CN,	CU,	CZ,	DE,	
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			CM,	GA,	GN,	ML,	MR,	NE,	SN,	TD,	TG								
	CA	2293	439			A1		1998	1217		CA	1998-	2293	439		1	9980	612	
	AU	9880	035			A		1998	1230		AU	1998-	8003	5		1	9980	612	
								2000											
	$z_{A}$	9805	140			A		1999	0107		$z_{A}$	1998-	5140			1	9980	612	
											EP	1998-	9280	10		1	9980	612	
	EP	1019	087			B1		2007	1121										
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IT,	LI,	LU,	NL,	SE,	MC,	PT,	
			ΙE,	FI,	CY														
		5016						2000	1222		NZ	1998-	5016	41		1	9980	612	
		2002						2002	0205		JΡ	1999-	5011	50		1:	9980	612	
		6814						2004	1109			2000-							
		1026						2008	0606		HK	2000-	1060	85		2	0000	926	
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ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT
OSC.G 5 THERE ARE 5 CAPLUS RECORDS THAT CITE THIS RECORD (8 CITINGS)
RE.CNT 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT